L1

## (FILE 'HOME' ENTERED AT 14:57:31 ON 27 JAN 2003)

FILE 'CAPLUS	' ENTERED AT 14:57:41 ON 27 JAN 2003
E	MASCAGNI PAOLO/IN,AU
132 S	E2-4
E	BOTTONI GIUSEPPE/IN, AU
12 S	E2-3
	71 OD 70

L2	12 S E2-3
L3	141 S L1 OR L2
L4	1777 S PAROXETINE
L5	2 S L3 AND L4
L6	22910 S CYCLODEXTRIN
L7	8 S L4 AND L6
L8	7 S L7 NOT L5

=> d ibib ab 1-2 L5 ANSWER 1 OF 2 CAPLUS COPYRIGHT 2003 ACS ACCESSION NUMBER: 2002:850956 CAPLUS DOCUMENT NUMBER: 137:316052 Pharmaceutical compositions comprising a TITLE: paroxetine salt and a polyhydroxylated INVENTOR(S): Mascagni, Paolo; Bottoni, Giuseppe PATENT ASSIGNEE(S): Italfarmaco S.p.A., Italy SOURCE: Ital. Appl., 27 pp. CODEN: ITXXCZ DOCUMENT TYPE: Patent Italian LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION: APPLICATION NO. DATE PATENT NO. KIND DATE -----IT 99MI1524 A1 20010112 IT 1999-MI1524 19990712 IT 1999-MI1524 PRIORITY APPLN. INFO.: 19990712 Title compns. are disclosed which are characterized by stability and absence of hygroscopicity and are prepd. by a process employing an aq. medium free of org. solvents. ANSWER 2 OF 2 CAPLUS COPYRIGHT 2003 ACS ACCESSION NUMBER: 2001:31497 CAPLUS DOCUMENT NUMBER: 134:105853 Preparation of complexes of paroxetine with TITLE: cyclodextrins or derivatives Mascagni, Paolo; Bottoni, Giuseppe INVENTOR(S): PATENT ASSIGNEE(S): Italfarmaco S.p.A., Italy SOURCE: PCT Int. Appl., 34 pp. CODEN: PIXXD2 DOCUMENT TYPE: Patent LANGUAGE: English FAMILY ACC. NUM. COUNT: PATENT INFORMATION: PATENT NO. KIND DATE APPLICATION NO. DATE ----------20010111 WO 2000-EP6121 20000630 WO 2001002393 A1 W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG IT 99MI1459 A1 20010102 IT 1999-MI1459 19990701 CA 2000-2341984 20000630 CA 2341984 AA 20010111 EP 1109806 A1 20010627 EP 2000-940418 20000630 R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, MC, IE, SI, LT, LV, FI, RO 20000630 BR 2000006838 Α 20010807 BR 2000-6838 PRIORITY APPLN. INFO.: IT 1999-MI1459 A 19990701 A 19991117 W 20000630 IT 1999-MI2406 WO 2000-EP6121 AΒ Complexes of paroxetine, as a free base or salt are prepd. with

AB Complexes of paroxetine, as a free base or salt are prepd. with a cyclodextrin or a cyclodextrin deriv. having a molar ratio between paroxetine and cyclodextrin ranging from 1:0.25 to 1:20, and these complexes are suitable for use in liq. and solid pharmaceutical compns. for oral and parenteral administration. Thus, a complex was prepd. from paroxetine and .beta.-cyclodextrin in a 1:1 ratio and the complex was characterized by NMR and thermal data. Tablets were prepd. from this complex and other excipients.

REFERENCE COUNT: 2 THERE ARE 2 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

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ANSWER 1 OF 7 CAPLUS COPYRIGHT 2003 ACS
ACCESSION NUMBER: 2002:695714 CAPLUS
DOCUMENT NUMBER:
                           137:222063
TITLE:
                           Serotonin reuptake inhibitor formulations
INVENTOR(S):
                           Chen, Chih-Ming; Li, Boyong; Cacace, Janice
PATENT ASSIGNEE(S):
                           Andrx Corporation, USA
                           PCT Int. Appl., 36 pp.
SOURCE .
                           CODEN: PIXXD2
DOCUMENT TYPE:
                           Patent
LANGUAGE:
                           English
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION: ~
     PATENT NO. KIND DATE
                                            APPLICATION NO. DATE
                                               -----
     WO 2002069888 A2 20020912
WO 2002069888 A3 20021227
                                              WO 2002-US4401 20020214
         W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
             CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR,
              LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT,
              RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZM, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR,
              BF, BJ, CF, CG, CI, CM, GA, GN, GQ, GW, ML, MR, NE, SN, TD, TG
                                           US 2001-785040 20010216
US 2001-785040 A 20010216
     US 2002156066
                      A1 20021024
PRIORITY APPLN. INFO.:
   A process for prepg. amorphous paroxetine-HCl or sertraline-HCl
     is provided, which comprises prepg. a soln. in which paroxetine
     -HCl or sertraline-HCl and a water-sol. polymer is dissolved in a
     co-solvent of a volatile org. solvent and water. Thus, granules were
     obtained from paroxetine-HCl 44.43, Povidone-K30 88.86, and
     Avicel PH-101 88.86 mg/tablet. The granules were blended with
     Cospovidone, microcryst. cellulose and Mg stearate to give a blend. This
     blend was compressed into tablets with a tablet wt. of 400 mg.
    ANSWER 2 OF 7 CAPLUS COPYRIGHT 2003 ACS
ACCESSION NUMBER:
                          2001:597801 CAPLUS
DOCUMENT NUMBER:
                           135:157705
TITLE:
                           Water dispersible formulation of paroxetine
INVENTOR (S):
                           Al-Ghazawi, Ahmad Khalaf Al-Deeb; Elder, David Philip;
                           Meneaud, Padma
PATENT ASSIGNEE(S):
                           Smithkline Beecham P.L.C., UK
                           PCT Int. Appl., 14 pp.
SOURCE:
                           CODEN: PIXXD2
DOCUMENT TYPE:
                           Patent
LANGUAGE:
                           English
FAMILY ACC. NUM. COUNT: 1
PATENT INFORMATION:
     PATENT NO.
                       KIND DATE
                                             APPLICATION NO. DATE
     WO 2001058449 A1 20010816
                                              -----
                        A1 20010816
                                              WO 2001-GB569
                                                                 20010209
         W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
              CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR,
             HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT,
             LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN,
             YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, TR, BF,
             BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                       A1 20021113
                                             EP 2001-904162 20010209
     EP 1255549
         {\tt R:} \quad {\tt AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT,} \\
     IE, SI, LT, LV, FI, RO, MK, CY, AL, TR
NO 2002003785 A 20020823 NO 2002-3
                                              NO 2002-3785
                                                                 20020809
PRIORITY APPLN. INFO.:
                                           GB 2000-3232
                                                          A 20000211
                                                             W 20010209
                                           WO 2001-GB569
     A water-dispersible formulation of paroxetine for immediate oral
     administration comprises a dry blend of paroxetine, a water-sol.
     dispersing agent, and a taste-masking agent, as a dispersible powder or
     molded into a tablet. For example, a water suspension contg.
     paroxetine, methacrylic acid copolymer, talc, and tri-Et citrate
     was spray dried. The spray dried material and polyvinylpyrrolidone,
     calcium carbonate, microcryst. cellulose, citric acid, flavor, sweetener,
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and Mg stearate were sieved, blended, and then compressed into tablets.

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THERE ARE 7 CITED REFERENCES AVAILABLE FOR THIS
REFERENCE COUNT:
                           7
                                  RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
L8 ANSWER 3 OF 7 CAPLUS COPYRIGHT 2003 ACS
ACCESSION NUMBER:
                           2001:300514 CAPLUS
DOCUMENT NUMBER:
                           134:331617
                           Oil-in-water emulsion compositions for polyfunctional
TITLE:
                           active ingredients
                           Chen, Feng-jing; Patel, Mahesh V.
INVENTOR(S):
PATENT ASSIGNEE(S):
                           Lipocine, Inc., USA
                           PCT Int. Appl., 82 pp.
SOURCE:
                           CODEN: PIXXD2
DOCUMENT TYPE:
                           Patent
                           English
LANGUAGE:
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
     PATENT NO.
                       KIND DATE
                                              APPLICATION NO. DATE
                              20010426
                                              WO 2000-US28835 20001018
     WO 2001028555
                        A1
         W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
              CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR,
              HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU,
              SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
          RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
              DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                                           US 1999-420159 19991018
US 1999-420159 A 19991018
                       A1 20020808
     US 2002107265
PRIORITY APPLN. INFO.:
   Pharmaceutical oil-in-water emulsions for delivery of polyfunctional
     active ingredients with improved loading capacity, enhanced stability, and
     reduced irritation and local toxicity are described. Emulsions include an
     aq. phase, an oil phase comprising a structured triglyceride, and an
     emulsifier. The structured triglyceride of the oil phase is substantially
     free of triglycerides having three medium chain (C6-C12) fatty acid
     moieties, or a combination of a long chain triglyceride and a
     polarity-enhancing polarity modifier. The present invention also provides
     methods of treating an animal with a polyfunctional active ingredient,
     using dosage forms of the pharmaceutical emulsions. For example, an
     emulsion was prepd., with cyclosporin A as the polyfunctional active
     ingredient dissolved in an oil phase including a structured triglyceride
     (Captex 810D) and a long chain triglyceride (safflower oil). The compn.
     contained (by wt.) cyclosporin A 1.0, Captex 810D 5.0, safflower oil 5.0,
     BHT 0.02, egg phospholipid 2.4, dimyristoylphosphatidyl glycerol 0.2,
     glycerol 2.25, EDTA 0.01, and water up to 100%, resp.
                                 THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS
REFERENCE COUNT:
                                  RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
    ANSWER 4 OF 7 CAPLUS COPYRIGHT 2003 ACS
ACCESSION NUMBER:
                           2001:136991 CAPLUS
DOCUMENT NUMBER:
                           134:198075
TITLE:
                           Triglyceride-free compositions and methods for
                           enhanced absorption of hydrophilic therapeutic agents
INVENTOR(S):
                           Patel, Mahesh V.; Chen, Feng-Jing
PATENT ASSIGNEE(S):
                           Lipocine, Inc., USA
                           PCT Int. Appl., 113 pp.
SOURCE:
                           CODEN: PIXXD2
DOCUMENT TYPE:
                           Patent
LANGUAGE:
                           English
FAMILY ACC. NUM. COUNT: 4
PATENT INFORMATION:
     PATENT NO.
                       KIND DATE
                                              APPLICATION NO. DATE
     WO 2001012155
                        A1
                              20010222
                                              WO 2000-US18807 20000710
         W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN,
              CR, CU, CZ, DE, DK, DM, DZ, EE, ES, FI, GB, GD, GE, GH, GM, HR,
             HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU,
             SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA, ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
         RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
             DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
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US 1999-375636 19990817 EP 2000-947184 20000710

B1

A1

20011030

20020605

US 6309663

EP 1210063

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R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO, MK, CY, AL
     US 2001024658 A1 20010927
                                                US 2000-751968
     US 6458383
                        B2 20021001
                                            US 1999-375636 A 19990817
WO 2000-US18807 W 20000710
PRIORITY APPLN. INFO.:
     The present invention relates to triglyceride-free pharmaceutical compns.
     pharmaceutical systems, and methods for enhanced absorption of hydrophilic
     therapeutic agents. The compns. and systems include an absorption
     enhancing carrier, where the carrier is formed from a combination of at
     least two surfactants, at least one of which is hydrophilic. A
     hydrophilic therapeutic agent can be incorporated into the compn., or can
     be co-administered with the compn. as part of a pharmaceutical system.
     The invention also provides methods of treatment with hydrophilic
     therapeutic agents using these compns. and systems. For example, when a compn. contg. Cremophor RH40 0.30, Arlacel 186 0.20, Na taurocholate 0.18,
     and propylene glycol 0.32 g, resp., was used, the relative absorption of
     PEG 4000 as a model macromol. drug was enhanced by 991%.
                                  THERE ARE 1 CITED REFERENCES AVAILABLE FOR THIS
REFERENCE COUNT:
                           1
                                  RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
    ANSWER 5 OF 7 CAPLUS COPYRIGHT 2003 ACS
                           2000:841959 CAPLUS
ACCESSION NUMBER:
DOCUMENT NUMBER:
                            134:21450
                            A pharmaceutical composition containing an active
TITLE:
                            agent in solid amorphous form
                            Chen, Jinling; Vilkov, Zalman
INVENTOR (S):
PATENT ASSIGNEE(S):
                            Purepac Pharmaceutical Co., USA
                            PCT Int. Appl., 38 pp.
SOURCE:
                            CODEN: PIXXD2
DOCUMENT TYPE:
                            Patent
LANGUAGE:
                            English
FAMILY ACC. NUM. COUNT:
PATENT INFORMATION:
      PATENT NO. KIND DATE
     PATENT NO.
                                                APPLICATION NO. DATE
     WO 2000071098 Al 20001130
                                               WO 2000-US14049 20000523
          W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR,
              \mathtt{CU},\ \mathtt{CZ},\ \mathtt{DE},\ \mathtt{DK},\ \mathtt{DM},\ \mathtt{DZ},\ \mathtt{EE},\ \mathtt{ES},\ \mathtt{FI},\ \mathtt{GB},\ \mathtt{GD},\ \mathtt{GE},\ \mathtt{GM},\ \mathtt{HR},\ \mathtt{HU},
              ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, PL, PT, RO, RU, SD,
              SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, UZ, VN, YU, ZA,
         ZW, AM, AZ, BY, KG, KZ, MD, RU, TJ, TM
RW: GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW, AT, BE, CH, CY,
              DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE, BF, BJ,
              CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG
                        A1 20020313
                                               EP 2000-936175 20000523
     EP 1185251
         R: AT, BE, CH, DE, DK, ES, FR, GB, GR, IT, LI, LU, NL, SE, MC, PT, IE, SI, LT, LV, FI, RO
PRIORITY APPLN. INFO.:
                                            US 1999-317448 A 19990524
WO 2000-US14049 W 20000523
     This invention relates to a pharmaceutical compn. and a process for
     producing a pharmaceutical compn. that contains an active agent in solid
     amorphous form wherein the amorphous form of the active agent is
     maintained. The active agents include paroxetine.cntdot.HCl
     (I), spironolactone, etodolac, and salts of diclofenac. I was dissolved
     in ethanol. The soln. was then mixed with complexing agent Crospovidone
     and co-solvent polyethylene glycol 300. After removing ethanol from the
     mixt., I in solid amorphous form was obtained.
REFERENCE COUNT:
                                  THERE ARE 5 CITED REFERENCES AVAILABLE FOR THIS
                           5
                                  RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT
    ANSWER 6 OF 7 CAPLUS COPYRIGHT 2003 ACS
ACCESSION NUMBER:
                           1999:377824 CAPLUS
DOCUMENT NUMBER:
                           131:164926
TITLE:
                           Separation of eleven central nervous system drugs by
                            capillary zone electrophoresis
AUTHOR (S):
                            Pucci, V.; Raggi, M.; Kenndler, E.
CORPORATE SOURCE:
                           Institute for Analytical Chemistry, University of
                           Vienna, Vienna, A 1090, Austria
SOURCE:
                           Journal of Chromatography, B: Biomedical Sciences and
                           Applications (1999), 728(2), 263-271
CODEN: JCBBEP; ISSN: 0378-4347
PUBLISHER:
                           Elsevier Science B.V.
DOCUMENT TYPE:
                           Journal
LANGUAGE:
                           English
   Several strategies to improve the sepn. of 11 central nervous system drugs
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(antipsychotics and antidepressants) with capillary zone electrophoresis were applied: the variation of the pH of the buffering background electrolyte, its ionic strength, addn. of inclusion-complex forming .beta.-cyclodextrin or polyvinylpyrrolidone (PVP), resp., as a replaceable, sol., polymeric pseudo-stationary phase. Best sepn. was achieved at pH 2.5 and 35 mmol/l ionic strength (phosphate buffer), with 0.5% (w/v) PVP. REFERENCE COUNT:

THERE ARE 24 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 7 OF 7 CAPLUS COPYRIGHT 2003 ACS ACCESSION NUMBER: 1999:233798 CAPLUS

130:272021

DOCUMENT NUMBER: TITLE:

Amorphous paroxetine composition Ronsen, Bruce; El-Rashidy, Ragab Pentech Pharmaceuticals, Inc., USA

PATENT ASSIGNEE(S):

PCT Int. Appl., 33 pp.

INVENTOR(S): SOURCE:

CODEN: PIXXD2

DOCUMENT TYPE:

Patent

LANGUAGE:

English

17

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND DATE	APPLICATION NO. DATE	
WO 9916440	A1 19990408	WO 1998-US20435 19980930	
W: CA, CN,	JP, KR, MX, NO		
RW: AT, BE,	CH, CY, DE, DK,	ES, FI, FR, GB, GR, IE, IT, LU, MC,	NL,
PT, SE			
CA 2304594	AA 19990408	CA 1998-2304594 19980930	
ZA 9808938	A 19991005	ZA 1998-8938 19980930	
EP 1019053	A1 20000719	EP 1998-951989 19980930	
R: AT, BE,	CH, DE, DK, ES,	FR, GB, GR, IT, LI, LU, NL, SE, MC,	PT,
IE, FI			
JP 2001517700	T2 20011009	JP 2000-513576 19980930	
PRIORITY APPLN. INFO	).:	US 1997-940058 A 19970930	

WO 1998-US20435 W 19980930 A free-flowing, amorphous paroxetine hydrochloride compn. suitable as a therapeutic agent for premature ejaculation can be prepd. by dissolving paroxetine free base in a hydrochloric acid-ethanol soln. followed by drying. The present compns. comprise amorphous paroxetine hydrochloride and at least one hydroxyl-bearing compd. In one preferred embodiment, the hydroxyl-bearing compd. is ethanol and the amt. of ethanol present in the amorphous product is in the range of 1-4 % based on paroxetine hydrochloride. The amorphous product is stable and substantially non-hygroscopic.

REFERENCE COUNT:

THERE ARE 17 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT